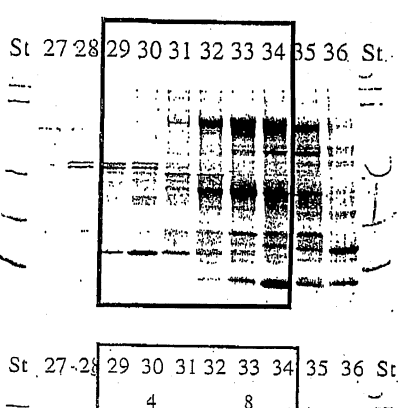


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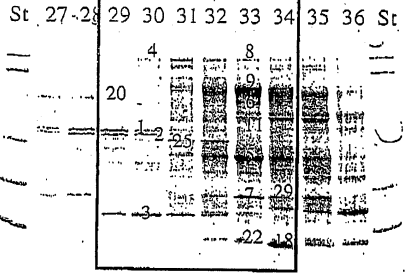


FIGURE # 3



Band No.	Identity
1	histone H1.c
2	histone H1.c
3	ribosomal protein RS20
4	similar to ribosomal protein LORP
5	BMP-3
6	α2 macroglobulin RAP & BMP-3
7	similar to ribosomal protein LORP
8	ВМР-3
9	BMP-3
11	ribosomal protein RL6 & BMP-3
18	TGF-β2/SPP24
20	Factor H
22	TGF-β2
25	BMP-3 & H1.x
29	BMP-3 & ribosomal protein RL32

FIGURE \$4

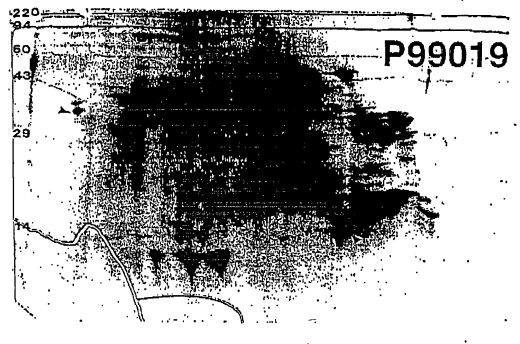
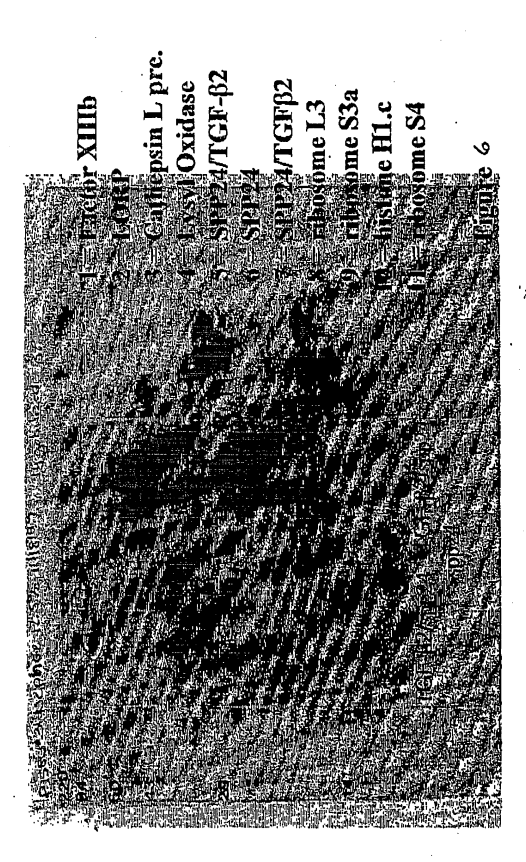
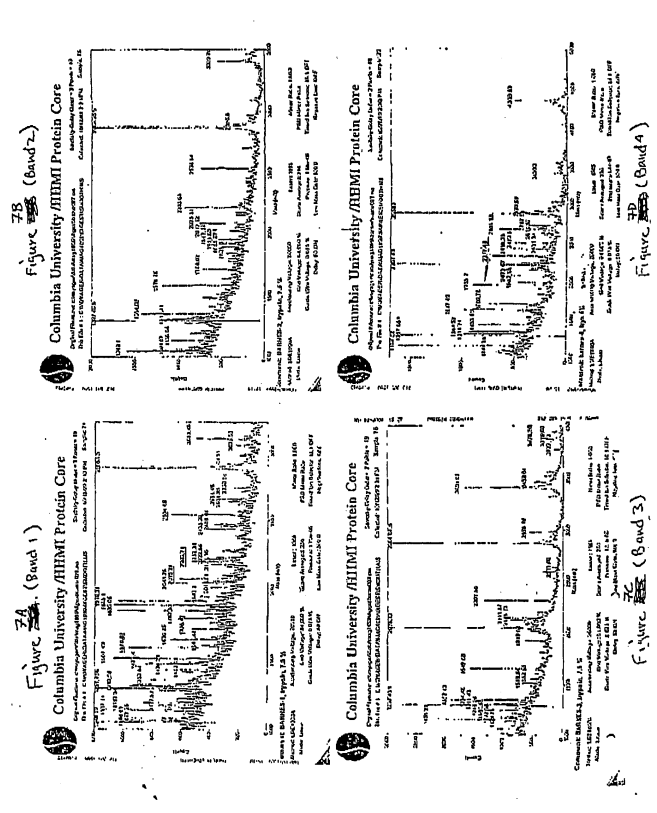
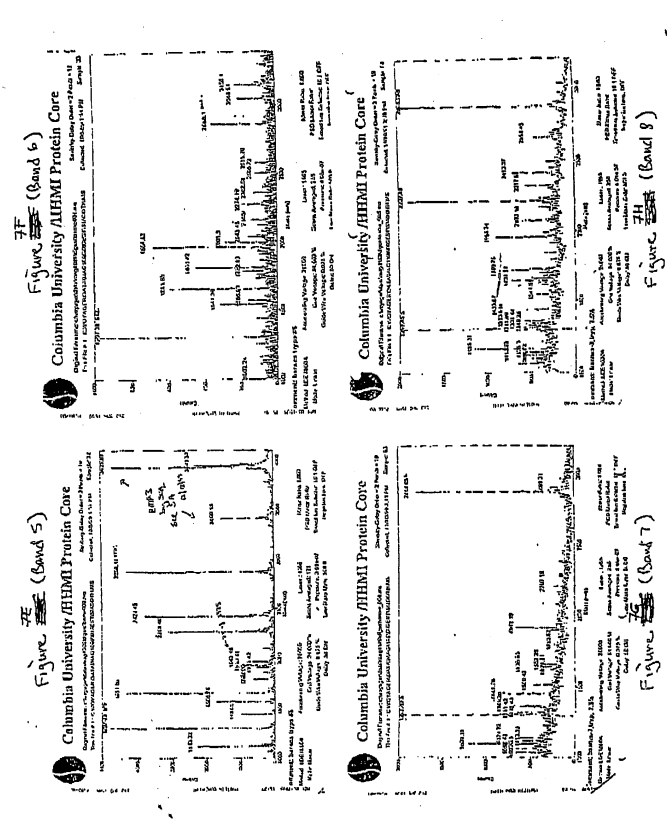


FIGURE #5

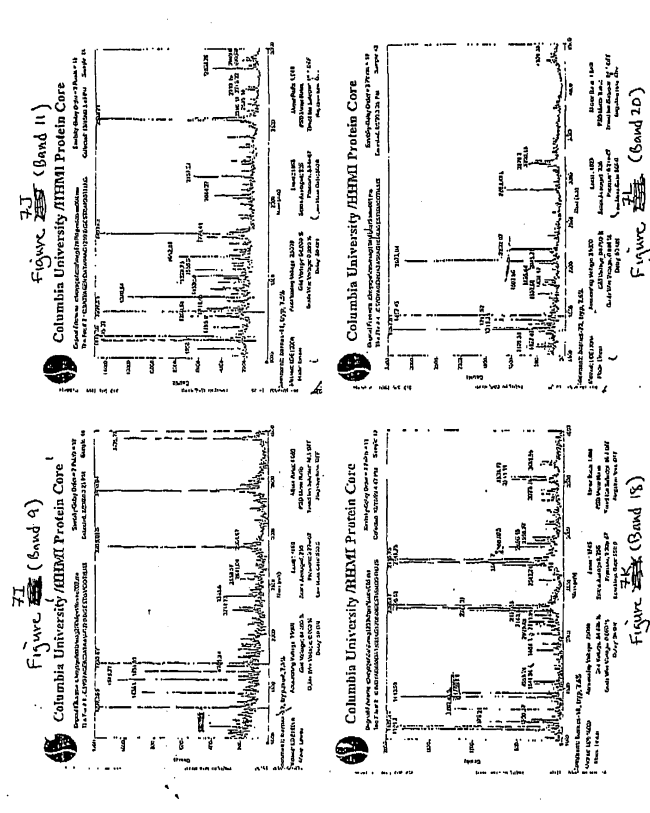




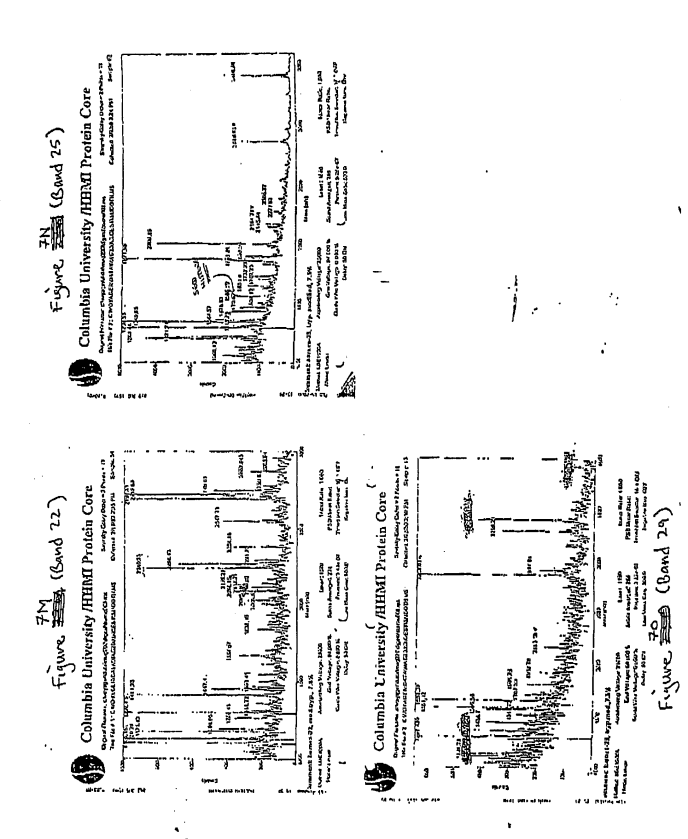


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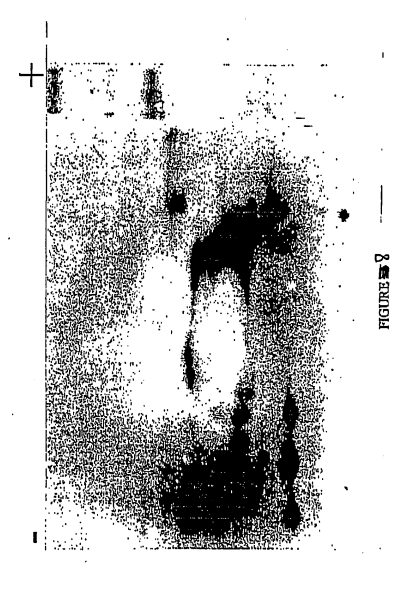


FIGURE # 98

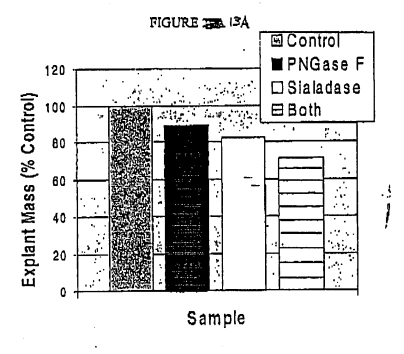
FIGURE ES 9C

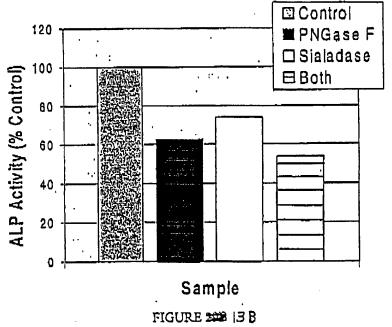
FIGURE SEP TD

FIGURE BY 9A



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Figure |4 Antibody Listing

Specificity	Antigen	Host	FCMC	Source	Catalog No.
		Species			•
TGF-b1 (burnan)	Protein	Rabbit	Polyclonal	Promega	G1221
TGF-b2 (furman)	Peptide	Rabbit	Polyclonal	Santa Cruz Biotechnology	06-38
TGF-b3 (hman)	Peptide	Kabbit	Polyclonal	Santa Cruz Biotechnology	sc-82
BMP-2 (laman)	Protein	Rabbit	Polyclonal	Austral Biologics	PA-513-9
BMP-3 (human)	Peptide	Chicken	Polyclonal	Research Genetics	NA
BMP-4 (human)	Peptide	Goat	Polyclonal	Santa Cruz Biotechaology	sc-6896
BMP-5 (human)	Peptido	Goat	Polyclonal	Santa Cruz Biotechnology	32-7405
BMP-6 (human)	Peptide	Mouse	Monoclonal	Novocustra Laboratories	NCL-BMP6
BMP-7 (human)	Peptide	Rabbit	Polycional	Research Genetics	NA
FGF-1 (human)	Peptide	Goat	Pulyclonal	Sarda Cruz Biotechnology	sc-1884
osteonectin (bovine)	Protein	Mouse	Monoclonal	ELISO	AON-1
osteocalcin (bovine)	Protein	Rabbit	Polycional	Accurate Chemicals	A761/R1H
serum albumin (bovine)	Protein	Rabbit	Polyclonal	Chemicon International	AB870
transferin (human)	Protein	Chicken	Polyclonal	Chenicon International	AB797
apo-Al lipoprotein	Protein	Goat	Polyclonal	Chemicon International	AB740
(Jannan)					

Figure 15A Identification of Proteins by Amino Acid Sequencing of Tryptic Fingments from 1D Gels

Band	Band Sample	Sequence Data	Best Database Match Match Identification	Match		Species	Acc. No.	AAs
_								
2	fx 49 (1579)	XLAAAGYDVEK	ALAAAGYDVEK	11/11	histone Hf.c	human .	87568 (NCBI)	65-75
м	(1346)	SLEKYCADLIR	SLEKVCADLIR	15/71	40s Ribosomal Profein S20	त्य	R3RTZ0 (PIR)	31-41
.	lx 65 ()	(V)VCGMLGFPSEAPV	VVCGNLGFPGEKRV 11/14	17/14	LORP	asnow	AAC95338 (NCBI)	213 228
\$	N terminal seq	STGVLPLQNNELPG	STGVILPLONNELPG 15/15	15/15	ви г -з	វេយាងព	4557371 (NCBI)	32, 35
	fx 72 (3925)	STGVLLPLONNELPGA EYQY	STGVILFLONNELPG 20/20 AEYQY		BMP-3	វិទិសា	4557371 (NCBI)	호 5 5 5 5 5 5 5 5 5 5 7 5 7 5 7 7 7 7 7
	fx 74 (3409)	STGVLLPLQ	STGVLPLO	6/6	8kP-3	turman	4557371 (NCBI)	290- 288
œ.	6x 55 (1566)	(s)атағке	SQTLQFDE	7/8	ย _เ ก3	វាធាតា	4557371 (NCBI)	35 ES
	fx 47	WAAF	no makh		m			
	N terminal seq	HAGKYSREKNT(P)A(P HGGKYSREKNOPKP) 11/1/4	HGGKYSREKNOPKP		a2-Macroglobulin Receptor Assoc. Pro.	វាមការមា	P30533 (Swiss-Prof)	31-46
	fx 57 (1438)	<u> saтгағре</u> а	sanafdea	6/6	ВІЛР-З	thumat	4557371 (NCBI)	354
_	tx 57 (1852)	SLICESNIFA	SLKPSNHA	8/8	E-dMB	ារពានកា	4557371 (NCBI)	410- 417
~	tx 51 (1093)	AALRPLVKP	AALRPLVKP	9/3	60s Ribosomal Protein L32	аѕпош	P17832 (Swiss-Prot)	6 -1
	fx 37 (90 [45]	A(H)I(Q)VERYV	AIVER	5/5	60s Ribosomal Protein 132	asnow	P17832 (Swiss-Prol)	108- 113
	fx 37 (ne MS)	A(H)J(Q)VERYV	НОЅВКҮУ	आ	60s Ribosomal Protein	əsnow	P17932 (Swiss-Prof)	22-28
8	fx 78 ()	XALF(G)AOLGXALGPI	no malch					
6 2	fx 56 (1587)	Satlafdeat	SQTLQFDEQT	10/10	BMP-3	nemuni.	P12645 (Swiss-Prof)	346. 355

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Figure 15B Identification of Proteins by Amino Acid Sequencing of Tryptic Fragments from 1D Gels.

			and Oatshace Match Match Identification	Matrh		Species	Acc. No.	AAs
Band	Band Sample	Sequence Data	מבשני הפוקדים			_		376
=	fx 55	Sature	Sance	8	BMP-3	רפשתייי	ASS 7.37 I	354
	(1311)		Т			T	Ī	5
	1x 47	VIATVTKPVGGDK	VLATVTKPVGGDK	13/13	60s Kibosoma) Proteia Lb Inuman	_	מיסקים מיסקים	27-10
	11772					T	(Synso-ring)	
_,	14 × 3	vWFA1	VFAL	414	60s Ribosomal Profein L6 [human			273
	17051						(Swiss-Prof)	9/2
		AVEOLOGYI B	AIPOLOGYI R	9/10	Sos Ribosomal Protein L6	human	Q02878	262-
	SPILL 111451	ייין ליינילטוניין					(Sviss-Prof)	13
<u>L</u> :								
			02020	0/0	TOE 82	human	P08112	303-
77.	2 28 2 28	ALDARYCFK	שויישורש)			(Swiss-Prot)	311
_	(1011)			;	101	1	DAR449	360
	(x 69 (no	GYNANFCAGACPYL	GYNANFCAGACHYL 14/14	14/14	71-451		rooms Swiss-Pmil	353
	Talet?						,	
	1% E6	VNSQSLSPY	WNSDSLSPY	6/6	SPP24	bovine	02/36/	42-5u
	(1411.71)						(SMSS-Prof)	
<u> </u> 2	EK 33	KAAKPSV(P)	KAAKPSVP	6/6	Histone Hf.x	human	JC4926 (PIR)	199-
<u>. </u>	(1470)							ş
٦						_		
					1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1			
,								

fx = fraction number (molecular weight of Gragment, as measured by SDS-PAGE)

Figur. 12 Identification of Proteins by Mass Spectrometry of Tryptic Fragments from 1D Gels

	•																			
Comments		15 MS peaks match with	Band 2				identification of stanted papelide confirmed by sequence analysis		15 MS peaks match with Band 1								12 MS peaks-match with Band 8	Į		
% Cover-	age	22					16			62							9	1	.•	
AAS		110-121		65-79	64-78	35-54	65-79	64-79	35-54	50-59	76-83	58-68	BB-89	B-21	5-21	BB-119	150-167		648-669	455-478
Mass Diff.	erace	09'0		0.16	0.58	-0.74	0.05	0.13	.0.20	0.36	-0.09	-0.16	0.55	720	-0.17	-0.85	-0.32		-0.28	0.47
Mass	Spec Database	1172.37		1579.71	1707.89	2012.32	1579.71	1707.89	2012.32	1129.40	1156.30	1334,62	1351,58	1517.77	1919.19	3404.87	1988.ZT	· .	2410.63	2610,10
Mass	Spec	1172.97		1579.87	1708.47	2011.58	1579.76	1708.02	2012.12	1129.76	1156.21	1334,46	1352,13	1518.04	1919.02	3404.02	1987.95	-	2410.35	2610.57
Acc. No.		87668	(MCBI)				87668 (NCBI)		•	R3RTZ0 (P.FC)			•				NP002309	90£		
Species Acc. No.	,	human					human			ख			,				buman			
Mass Spet	Profile	4 peaks	match with histone H1.c				3 peaks malch with histone H1.c			7 peaks match with nbosome S20							3 peaks match with	Lysyl Oddase RP		
Band		-	•				2			က							4			

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Figure. 16B Identification of Proteins by Mass Spectrometry of Tryptic Frugments from 1D Gels

																			_
Comments		% coverage calculation is relative to the mature BMP-3, 183 AAS (290-472)			-			temets lo noiteatimati	peplide confirmed by sequence analysis								% coverage calculation is relative to the mature BMP-3, 163 AAS (290-472)		
% Cover-	age	48									17						15		
AAS		361-368	346-357	345-55	410-424	346-350	374-392	373-392		200-318*	283-290				129-150	257-282	346-357	410-424	
Mass Diff-	erence	0.01	-0.05	0.00	-0.05	70.0	-0.17	-0.36	· · · · · · · · · · · · · · · · · · ·	1.38	0.09				0.15	-0.01	0,18	-0.03	
Mass	Spec Database	1113.31	1438.58	1585.76	1651.91	1794.02	2268.63	2424.81		3407.77	1002.15				238243	3048.52	1566.75	1651.91	
Mass	Spec	1113.32	1438.53	1566.76	1651.86	1794.09	2258.46	2424.45		3409.15	1002,24		~.		2362,58	3048.51	1565.93	1651,88	
Acc. No.		4557371 (NCBI)									P30533	(Swiss-	Prot)				4557371 (NCBI)		
Species		human									human						human		
Mass Sper		9 peaks match with AMP-3									3 peaks	match with	4.5	Macroglobuli n RAP	2		2 peaks match with BMP-3		
Band		s									9								

Figure 16. Libertification of Proteins by Mass Spectrometry of Tryptic Fragments from ID Gels

Comments		•		,				% coverage catculation is	3, 183 AAS (290-472)					12 MS peaks match with Band 4		% coverage calculation is	relative to the mature BMP- 3, 183 AAS (290-472)				-	
% Corer	age		2					7						૯		36					1	
AAs			67-75		1-10	65-74	19-28	102-111		351-368	190-200	410-424	346-360	648-869		361-368		346-357	345-357	410424	41-66	290-318
Mass Dill-	erence		0.08		-0.09	0.44	0.12	0.22		0.08	-0.32	0.37	-0.40	-0.26	,	1710		0.02	0.01	0.30	0.48	1.17
М	Sper	Database	1033.17		1083.40	1134.28	1449.68	1060.20		1113.31	1360.58	1651.91	1794.02	2410.63		4443 21		1438.58	1566.76	1651.61	2901.19	3407.77
Mass		Darla	1033.25		1093.31	1134.72	1448.78	1050.42		1113.39	1380,28	1652.28	1793.62	2410.37		4449 44	5	1438.60	1586.77	1651.91	2901.67	3408.94
Acc. No.	•••		P17932	Prof)	- Take MA	,		4557371	(NCBI)				. •	NP002309	Prod)	150031	(NCBI)	سنيد				
Species			mouse			-		human						human								
Mass Spec	Profile	-	4 peaks	match with ibosome	757			5 peaks	match with BMP-3					1 peak	Lysyl Oxidase RP		match with					
Band	· ·		7						- 11 - 12 -					8		ŀ	n					

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Figure 16D Identification of Proteins by Mass Spectrometry of Tryptic Fragments from 1D Gels

				•																					_
Comments		% coverage calculation is making RMP.	3, 183 AAS (290-472)								•														
% Cover-	age	48						16							29					ଚ୍ଚ		1	•		
AAs	·	361-358		410-424	348-360	373-392	290-318	114-122		141-155	10-20		262-271	260-271	303-311		400-409	312-328	340-382	42-53		113-124	98-98	62-77	
Mass Diff-	etence	60.0-		-0.18	-0.44	15.0-	0.57	0.15		D.02	0.03		0.01	0.00	-0.06		-0.16	-0.23	-0.21	19'0-		-0.06	10 0	50.0	-0.10
Mass	Spec Database	1113.31		1651.91	1794.02	2424.81	3407.77	114023		1526.86	1059.12		1145,35	1366.69	1101,26		1175.42	2240.80	1631692	1411,60		1447.65	1540.60	1889.05	2268.57
Mass	Spec	1113.23		1651.73	1793.58	242424	3408.34	1140.38		1526.88	1059.15	•	1145.38	1386.74	1101.20		1175.26	7240.37	2691.70	1410.93		1447.59	1540.94	1869.10	2268.47
Acc. No.		4557371	(NCBI)					Q02878	(Swiss- Prof)		P47911	(Swiss-	?		P08172	Prof)				796720	(Swiss-				
Species	•	human						human			Mouse				human					bovine				•	
Mass Spec	Profile	5 peaks	malch with BMP-3					5 peaks	match with						4 peaks	metch with TGF-02				5 peaks	match with		-		
Band		=													92										

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figure. ICE Identification of Proteins by Mass Spectrometry of Tryptic Fragments from 1D Gels

																	_			
Comments													1		% coverage calculation is	3, 183 AAS (290-472)				
% Cover-	age	83					11			14					34	•	•			
AAS		303-311	400-409	312-347	312-328	340-362	42-53		113-124	48-57	107-118	107-119	48-58	43-57	102-111		346-357	345-357	410-424	290-318
Mass Diff-	erence	-0.11	-0.29	-0.26	-0.35	-0.30	-0.37		-0.25	0.05	-0.64	-0.67	-0.02	-0.74	0.23		0.25	0.15	-0.11	1,09
Mass	Spec Database	1103.26	1175.42	2084.42	2240.60	2691.91	1411.60		1447.65	1208.40	1222.35	1350.52	1384.59	1732.97	1060.20		1438.58	1569.76	1551.91	3407.77
Mass	Spec Data	1101.15	1175.13	2084,18	2240.25	2691.61	1411.23		1447.40	1208.46	1221.71	1349.85	1384.57	1732.23	1060.43		1438.83	1566.92	1651.80	3408.86
Acc. No.		Po8112 (Swiss- Prof)					027967	(Swiss- Prof)		JC4828 (PIR)					4557371	(NCBI)				
Species	•	human					bovina			human					ћитав			-		
Mass Spec		5 peaks match with TGF-82					2 peaks	match with SPP24		5 peaks snaich with					5 peaks	match with BIAP-3				
Band		22								25										

Figure 16F Identification of Proteins by Mass Spectrometry of Tryptic Vraguents from 1D Gels

Соптепts	% coverage calculation is relative to the mature BMP-3, 183 AAS (290-472)			
% Cover- age	27	~=		
AAs	-0.09 381-368		345-357	
Mass Mass Diff- Spec erence atabase	-0.09	0.12	0.11	127
Mass Sper Database	1113.31	1438.5B	1568.75	3407.77
Mass Spec Data	1113.22	1438.70	1566.66	3408.04
Acc. No.	4557371 (NCBI)			
Species	human			
Band Mass Spec Species Acc. No.	4 peaks match with RMP-3			
Band	23			

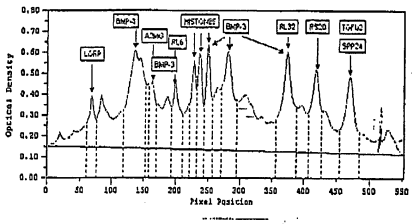


FIGURE 17A



FIGURE 178



Figure 18 Quantitation of Identified BP proteins

Identified Protein	Percentage of Total Protein
LORP	2
вмр-з	11
BMP - 3 & A2-MG	3
RL6 & BMP-3	4
Histone	3 - ,
Histone	3
Histone & BMP-3	4
ВМР-3	8
RL32 & BMP-3	8
R\$20	5
SPP24 & TGF-β2	6
Total	58%

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Figure PA Identification of Proteinsby Mass Spectrometry of Fragments from 2D Gels

								_				_			_,	_
Comments			•		peptide match	sequence analysis										
*	Сометаде		60				C)		41		_	_				
AAs			472-487	368-382			241-253	0.26 648-669	105-116	58-70	21-33	301-314	310-334	274-295	239-261	0.41 135-154
		Diff	-0.13	0.51	NA		-0.31	0.26	0,46	0.14	0.36	1.06	0.71	1.40	08'0	
		Dafabase	1837.14	1921 14	MA		1609.86	2410,63	1406.80	1546.70	1660.60	1680.80	1834.60	2351.50	2380.70	2725.10
MS	Peaks	Dada	1837,01	1921 65	2679.51		1609.57	2410.89	1467.26	1546.64	(661.16	1581.85	1834.71	2352.90	2381,50	2721.51
Acc. No.			P05160 (Swiss- Prot)				NP002309 (Swiss- 1609,57 Prof)	•	P25975 (Swiss- Prot)		•					
Species			Human				Human		Bovine							
Mass Spec			2 peaks match with	Factor XIIIb			2 peaks match		8 peaks match with Cethepsin	L Precursor			="			,
Dioest			Lysc				Trypsin		Lysc							
Spol	<u>.</u>		-				7		m							

Figure 19C Identification of Proteinsby Mass Spectrometry of Fragments from 2D Gets

					-т			 -	 -Y				-7	~	7		7	٦		~	~
Comments																					
*	Сочегаде		42				10			37				_							
AAS			26-31	32-37	98-107	1-25	42-60			348-355	10-18	286-296	249-260	103-114	103-115	65-78	30-49	177-197	200-223	70-98	-0.25 198-223
		Diff	-0.34	-0.25	-0.31	1.44	0.26			0.25	0.08	0.22	0.02	0.17	0.04	-0.16	-0.12	0.34	0.27	-0.49	
		Database	774.90	809.84	1175,43	3166.66	2187.51			917.14	984.15	1192.40	1380.65	1464.63	1620.82	1779.00	2238.55	2325.65	2561.04	2898.43	2948.35
SM	Peaks	Data	774.56	B09.69	1175.12	3168.10	2187.77			917,39	984.23	1192.62	1389.87	1464.60	1620.86	1778.64	223B.43	2325.39	2661.31	2897.94	2946.10
Acc. No.			P21214 (Swiss- Prol)				027967 (Swiss-	Prod		P39872 (Swiss- Prof)								•			
Species			Bovine		•		Bowine			Bovina											
Mass Spec			4 peaks malch		-		t peak	matches with SPP24	i i	Trypsin 12 peaks match Bovine	3									اشديت	
Dinest			Lys.C							Trypsin											
Sport			7																		

Figure #D Ideatification of Proteinsby Mass Spectrometry of Firzgments from 2D Gels

Comments																					
*	Coverage		28							23				1 23				_			
AAs			19-26	152-161	151-161	174-185	94-108	199-212	65-81	34-46	65-79	64-79	1-21	230-239	134-141	230-241	198-210	37-48	221-239	17-99	
		Diff	-0.05	-0.02	0.13	Q DO	-0.10	-0.09	-0.04	0.19	-D.01	-024	-0.38	0.10	00'0	0.42	0.12	-0.23	-0.24	-0.10	
		Database	920.10	1218.31	1348.49	1516,69	1593.82	1720.00	1953,16	1327.58	1579.71	1707.89	2147.53	1 (68.38	1216.39	1353.61	1507.69	1557.98	2140.58	2591.90	
MS	Peaks	Data	920.05	1218.20	1346.62	1516.69	1593.72	1715.91	1953,12	1327.75	1579 70	1707,65	2147.17	1168.48	1216.39	1354.03	1507.81	1557.75	2140.34	2591.80	
Acc. No.			P97351 (Swiss- Pro!)							87668 (NCBI)				P12750 (Swiss- Prot)							
Soeries			Mouse							Human				Human							
Mass Spec			7 peaks match with nbosome								71.C			Trypsin 6 peaks match Human with nbosome	ħ						
Dinect	15		Trypsin							Trypsin				Trypsin			<u> </u>				
Cont	וחתר		60					_		5				=	_						